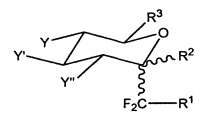
Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

1 (original). A gem-difluorinated compound of formula:



wherein

 ${\ensuremath{\mathsf{R}}}^1$ is a group comprising an alkyl chain substituted with at least one amine, amide, or acid function,

 ${\ensuremath{\mbox{R}}}^2$ is a hydrogen atom H or a free or protected alcohol function,

 R^3 is notably an H, CH_3 , CH_2OH , CH_2-OGP group wherein GP is a protective group such as an alkyl, benzyl (Bn), trimethylsilyl (TMS), tert-butyl-dimethylsilyl (TBDMS), tert-butyldiphenylsilyl (TBDPS), acetate (Ac)...,

Y, Y', Y" are independent groups wherein Y, Y', Y" = H, OR, N₃, NR'R", SR'" ... with R = H, Bn, Ac, TMS, TBDMS, TBDPS, ..., R', R" = H, alkyl, allyl, Bn, tosylate (Ts), C(=0) - alkyl, C(=0) -Bn, ..., R''' = H, alkyl, Ac.

2 (currently amended). The compound according to claim 1, characterized in that it comprises comprising a Cglycoside of general formula:

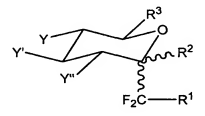
wherein R^5 and R^6 = H or a group either functionalized or not such as a functionalized carbon chain bearing i.a. an amine, amino acid, aminoester function, a peptide chain, a protein, a carbohydrate, a steroid, or a triterpene, an alkaloid, a lignane, or compounds of pharmacological interest.

3 (currently amended). The compound according to claim 1, characterized in that it comprises comprising a glycoconjugated compound of general formula:

$$R^3$$
 Y''
 Y'''
 R^2
 R^3
 R^2
 R^3
 R^2
 R^2
 R^3
 R^2
 R^2
 R^3
 R^3

wherein R^5 , R^6 , R^7 and R^9 = H or a group either functionalized or not, such as a functionalized carbon chain bearing i.a. an amine, amino acid, aminoester function, a peptide chain, a protein, a carbohydrate, a steroid, or a triterpene, an alkaloid, a lignane, or compounds of pharmacological interest.

4 (currently amended). A method for preparing a gemdifluorinated compound of formula:



wherein

 ${\ensuremath{\mathsf{R}}}^1$ is a group comprising an alkyl chain substituted with at least one amine, or amide function,

 ${\ensuremath{\mbox{R}}}^2$ is a hydrogen atom H or a free or protected alcohol function,

 R^3 is notably an H, CH_3 , CH_2OH , CH_2 -OGP group wherein GP is a protective group such as an alkyl, benzyl (Bn), trimethylsilyl (TMS), tert-butyl-dimethylsilyl (TBDMS), tert-butyldiphenylsilyl (TBDPS), acetate (Ac)...,

Y, Y', Y" are independent groups wherein Y, Y', Y" = H, OR, N_3 , NR'R", SR'" ... with R = H, Bn, Ac, TMS, TBDMS, TBDPS, ..., R', R'' = H, alkyl, allyl, Bn, tosylate (Ts), C(=O) - alkyl, C(=O)-Bn, ...,

R''' = H, alkyl, Ac,

characterized in that it comprises said method comprising a reaction between a lactone and a halogenated derivative of general formula $XCF_2CO_2R^8$, wherein X is a halogen, in the presence of zinc, or of a lanthanide derivative and R^8 = alkyl, aryl...

5 (currently amended). The method according to claim 4, characterized in that wherein said lanthanide derivative is samarium diiodide.

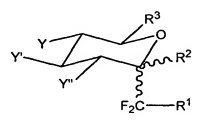
6 (currently amended). The method according to claim 4, characterized in that wherein said sugar derivative is obtained in one or more steps from a corresponding commercially available sugar.

7 (currently amended). The method according to claim 4, characterized in that <u>wherein</u> said reaction is followed by a deoxygenation.

8 (currently amended). The method according to claim 4, $\frac{1}{2}$ characterized in that wherein the R⁸ group comprises an ester function which is reduced to alcohol.

9 (currently amended). The method according to claim 4, characterized in that <u>wherein</u> the R⁸ group comprises an ester function which is either reduced to alcohol then oxidized into an aldehyde or hemiacetal, or directly reduced into aldehyde.

10 (currently amended). A method for preparing a gemdifluorinated compound of formula:



wherein

 $R^1 = -C(=0) - NR^5 R^6$, wherein R^5 and $R^6 = H$ or a group either functionalized or not, such as a functionalized carbon chain bearing i.a. an amine, amino acid, aminoester function, a peptide chain, a protein, a carbohydrate, a steroid, or a triterpene, an alkaloid, a lignane, or compounds of pharmacological interest,

 ${\ensuremath{\mbox{R}}}^2$ is a hydrogen atom H or a free or protected alcohol function,

R³ is an H, CH₃, CH₂OH, CH₂-OGP group wherein GP is a protective group such as an alkyl, benzyl (Bn), trimethylsilyl (TMS), tert-butyl-dimethylsilyl (TBDMS), tert-butyldiphenylsilyl (TBDPS), acetate (Ac)...,

Y, Y', Y" are independent groups

wherein Y, Y', Y" = H, OR, N_3 , NR'R", SR'" ...

with R = H, Bn, Ac, TMS, TBDMS, TBDPS, ...,

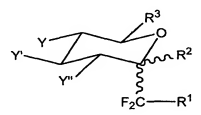
R', R'' = H, alkyl, allyl, Bn, tosylate (Ts), C(=O) -

alkyl, C(=0)-Bn, ...,

R''' = H, alkyl, Ac,

characterized in that it comprises said method comprising a Ugi reaction with an amine, an aldehyde and an isonitrile.

11 (currently amended). A method for preparing a gemdifluorinated compound of formula:



wherein

 $R^1 = -C(=0) - NR^5 R^6$, wherein R^5 and $R^6 = H$ or a group either functionalized or not, such as a functionalized carbon chain bearing i.a. an amine, amino acid, aminoester function, a peptide chain, a protein, a carbohydrate, a steroid, or a triterpene, an alkaloid, a lignane, or compounds of pharmacological interest,

 ${\ensuremath{\mbox{R}}}^2$ is a hydrogen atom H or a free or protected alcohol function,

R³ is an H, CH₃, CH₂OH, CH₂-OGP group wherein GP is a protective group such as an alkyl, benzyl (Bn), trimethylsilyl (TMS), tert-butyl-dimethylsilyl (TBDMS), tert-butyldiphenylsilyl (TBDPS), acetate (Ac)...,

Y, Y', Y" are independent groups
wherein Y, Y', Y" = H, OR, N₃, NR'R", SR'" ...
 with R = H, Bn, Ac, TMS, TBDMS, TBDPS, ...,
 R', R" = H, alkyl, allyl, Bn, tosylate (Ts), C(=0) alkyl, C(=0) -Bn, ...,
 R'" = H, alkyl, Ac,

characterized in that it comprises said method comprising a coupling reaction of a sugar derivative with an amine.

12 (currently amended). A composition, characterized in that it comprises comprising at least one compound according to claims 1 to 3 claim 1 or one of its derivatives or one of its salts obtained by addition to a pharmaceutically acceptable organic or mineral acid.

13 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing antitumoral drugs.

- 14 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing antiviral drugs.
- 15 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing hypoglycemic drugs.
- 16 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing compounds for immunology.
- 17 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing anti-inflammatory compounds.
- 18 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing compounds for cosmetology.
- 19 (currently amended). The use of a gem-difluorinated compound according to any of claims 1 to 3 claim 1, for preparing glycopeptide analogs of antifreeze molecules.